

JAN 6 2006

Attachment 1 - 510 (k) SUMMARY

510(k) summary for NDR+ – DIVA-D

Identification

Applicant	NICAL SPA		
	43 Via Soffredini		
	Milan (MI), ITALY 20126		
	Registration Number: 3003314115		
Contact Person	Mr. Roberto Niccolucci - President		
Telephone (applicant)	+39 022571110		
Designated Agent	Mr. Gerald Silverman		
in the US	71 Rose Street		
	Hasting On Hudson, NY 10706		
	Phone: +1 914 674 1085		
Manufacturing site	NICAL SPA		
	43 Via Soffredini		
	Milan (MI), ITALY 20126		
	Registration Number: 3003314115		

Trade name: NDR+/DIVA-D

Common name: Digital image acquisition system

Classification:

The equipment is classified as a class II

CFR21- 892.2050: Picture archiving and communication system

CFR21 - 892.1650: Image Intensified Fluoroscopic X-ray system.

Substantial equivalent device: the NDR+/DIVA-D is defined as Substantially Equivalent (SE) to the INFIMED Platinum ONE digital image acquisition system (certified under the name ORION) (K012490).

The following table compares the NDR+/DIVA-D and the predicate device



FEATURES	NICAL (NDR+/DIVA D)	INFIMED (Platinum ONE)
Intended use	The NICAL NDR+ is a digital image acquisition system to be used in conjunction with an image Intensifier during radiography or fluoroscopy x-ray examination to capture images by a camera, digitalize the image, review images and format images according to DICOM 3.0 protocol to be sent through network connection	The INFIMED Orion Fluoroscopic imaging system is a high resolution, digital imaging system designed for digital videography. It is intended to replace conventional film techniques in multipurpose or dedicated applications where general fluoroscopy, interventional fluoroscopy or angiography or cardiac imaging procedures are performed. The Orion system allows the operator to view and enhance 1000 line fluoroscopy. High resolution digital spot images (1024x1024) may be acquired at single or rapid acquisition rates. Images may be viewed and enhanced enabling the operator to bring out diagnostic details difficult or impossible to see using conventional imaging techniques. The Orion system enables the operator to hardcopy image with a laser printer or send images over a network. The major system components include: a fluoroscopic TV camera, monitors, and an image processor.
	GGD 1024 1024 121 iv	000
Image Acquisition Speed Acquisition	CCD camera 1024x1024 12 bit 1024x1024 at 25 fps for FLUORO and all radiographic exams	CCD camera 1024x1024 12 bit Up to 30 fps for FLUORO acquisition and up to 15 fps for spot acquisition
Edge Enhancer	Completely hardware in real time	Software package
Start Up System	Automatic hardware and software test for each ignition	Automatic system calibration and remote system diagnostics
Noise Reduction	Dynamic recursive filter which offering an automatic noise/blurring optimization effect it gives zero persistence	Standard recursive filter
Image Storage	Up to 36000 frames with matrix 1024x1024. All the images are stored directly from the CCD camera head without any kind of edge enhancement and digital compression.	NA
Post Processing Working	All the images stored in the system can be elaborated in order to modify the windowing, the gray scale, the zoom, the annotation package, the edge enhancement and the noise reduction	Brightness/contrast polarity enhancement. Measurement and annotation package

510(K) Summary for NDR+/DIVΛ-D



DSA Application	Real time mask subtraction, road	Real time mask subtraction, road
	mapping, land marking, pixel shift	mapping, land marking, percent
	function, auto loop replay, injector	Stenosis vascular package, auto loop
	synchronization, automatic or manual	replay, injector synchronization.
	possibility to change the mask, maxop	
	function and stepping angio	
Multi Image	Thumbnail image viewing with the	
	possibility to choose the first image for	Thumbnail image viewing
	every run or the 50% of the run.	
DICOM Functions	Full DICOM 3.0 integrated	Full DICOM 3.0 integrated
Laser Interface	When DICOM network is not	
	available in the site installation, it is	
	possible to send the images to a laser	NA
	printer using the protocol P831 or	
	P859	
Interface Commands	Flat keyboard with waterproof surface	Standard pc keyboard + standard pc
		mouse

Indication for use.

The indication for use of the NDR+/DIVA-D is: DIGITAL IMAGE ACQUISITION SYSTEM TO BE USED IN CONJUCTION WITH AN IMAGE INTESIFIER DURING RADIOGRAPHY OR FLUOROSCOPY X-RAY EXAMINATION TO CAPTURE IMAGES BY A CAMERA, DIGITALIZE THE IMAGE, REVIEW IMAGES AND FORMAT IMAGES ACCORDING TO DICOM PROTOCOL TO BE SENT THROUGH NETWORK CONNECTION



SFP - 6 2006

Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

NICAL SPA % Mr. Gerald Silverman Designated Agent 71 Rose Street HASTING ON HUDSON NY 10706

Re: K053029

Trade/Device Name: NDR+/DIVA-D Regulation Number: 21 CFR §892.1650

Regulation Name: Image-intensified fluoroscopic x-ray system

Product Code: MQB

Regulation Number: 21 CFR §892.2050

Regulation Name: Picture archiving and communications system

Product Code: LLZ
Regulatory Class: II
Dated: October 27, 2005

Received: November 25, 2005

Dear Mr. Silverman:

This letter corrects our substantially equivalent letter of January 6, 2006.

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (sections 531-542 of the Act); 21 CFR 1000-1050.

This letter will allow you to continue marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (240) 376-3150 or at its Internet address http://www.fda.gov/cdrh/dsma/dsmamain.html

Sincerely yours,

Nancy C. Brogdon

Director, Division of Reproductive,

Abdominal, and Radiological Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure

cc: Mr. Roberto Daglio General Manager Operations Villa Sistemi Medical via Azalee 3 20090 Buccinasco-Milan ITALY



6.1. Indication for use Statement

510(k) Number: 1/053029

Device Name: NDR+ or DIVA-D

The indication for use of the NDR+/DIVA-D is: DIGITAL IMAGE ACQUISITION SYSTEM TO BE USED IN CONJUCTION WITH AN IMAGE INTESIFIER DURING RADIOGRAPHY OR FLUOROSCOPY X-RAY EXAMINATION TO CAPTURE IMAGES BY A CAMERA, DIGITALIZE THE IMAGE, REVIEW IMAGES AND FORMAT IMAGES ACCORDING TO DICOM PROTOCOL TO BE SENT THROUGH NETWORK CONNECTION

(Division Sign-Off)
Division of Reproductive, Abdominal, and Radiological Devices 510(k) Number

Prescription Use